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Localization and Processing of the Amyloid- β Protein Precursor in Mitochondria-Associated Membranes

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Abstract. Alteration of mitochondria-associated membranes (MAMs) has been proposed to contribute to the pathogenesis of Alzheimer's disease (AD). We studied herein the subcellular distribution, the processing, and the protein interactome of the amyloid- β protein precursor (A β PP) and its proteolytic products in MAMs. We reveal that A β PP and its catabolites are present in MAMs in cellular models overexpressing wild type A β PP or A β PP harboring the double Swedish or London familial AD mutations, and in brains of transgenic mice model of AD. Furthermore, we evidenced that both β - and γ -secretases are present and harbor A β PP processing activities in MAMs. Interestingly, cells overexpressing APP_{swe} show increased ER-mitochondria contact sites. We also document increased neutral lipid accumulation linked to A β production and reversed by inhibiting β - or γ -secretases. Using a proteomic approach, we show that A β PP and its catabolites interact with key proteins of MAMs controlling mitochondria and ER functions. These data highlight the role of A β PP processing and proteomic interactome in MAMs deregulation taking place in AD.

Keywords: Alzheimer disease, amyloid- β protein precursor, lipids, mitochondria associated membranes, proteomic

INTRODUCTION

Alzheimer's disease (AD) is a devastating neurodegenerative pathology characterized by the

progressive decline of cognitive functions. The accumulation of intracellular aggregates of tau protein in the neurofibrillary tangles and of extracellular aggregates of a set of polypeptides called amyloid- β peptides (A β) in the senile plaques [1] are the major histological hallmarks of AD. A β peptides are generated from a large transmembrane protein, the amyloid- β protein precursor (A β PP) through its sequential cleavage by the β -secretase generating the

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C-terminal fragment (CTF) of 99 amino acids (C99), and by the γ -secretase releasing A β peptide, and the APP intracellular domain (AICD) [2]. Cleavage of A β PP by α -secretase precludes the formation of A β peptides and leads to the generation of another CTF of 83 amino acids (C83) [3, 4].

A β peptides are generated extracellularly, but also within intracellular organelles such as the endoplasmic reticulum (ER) and mitochondria. A β PP is folded and modified in the ER and transported through the Golgi complex to the plasma membrane. It was proposed that A β is generated in the ER lumen as a result of deficits in axonal transport [5]. A β oligomers (considered to be the most toxic forms of A β) accumulate in the ER in hippocampal neurons and thereby, likely lead to cell death in transgenic mice expressing A β PP mutant [6]. Association of A β PP with mitochondria was also reported by several groups [7–9]. A β was shown to accumulate into mitochondria prior to extracellular plaques and in a time-dependent manner, leading to mitochondrial damage [10–12]. In addition, different components of the γ -secretase complex have been located in different subcellular compartments, including the ER and mitochondria [9, 13–16].

Physical association between the ER and mitochondria, also known as mitochondria-associated membranes (MAMs), play important roles in various cellular housekeeping functions such as phospholipids-, glucose-, cholesterol-, and fatty-acid- metabolism, as well as calcium signaling [17, 18]. The hypothesis stating that AD could be a disorder linked to MAMs alterations started to emerge only in the last 6 years [19–21]. In this study, we show that A β PP and its catabolites accumulate in MAMs and increase ER-mitochondria contact sites. We also evidence that both β -secretase and γ -secretase complex are present and may harbor enzymatic activities in MAMs. Of most importance, we document increased neutral lipid accumulation in cellular models of AD, which was reversed by inhibiting β - or γ -secretases. A proteomic approach revealed the protein interactome of A β PP and its catabolites with key proteins of MAMs controlling mitochondria and ER functions.

MATERIAL AND METHODS

Cells

Human SH-SY5Y neuroblastoma cells (CRL-2266, ATCC) stably expressing pcDNA3.1 (control), wild type human A β PP (APPwt), or human A β PP

harboring the double Swedish mutations (APP^{swe}: APPKM670/671NL) constructs were generated as already described [22]. Polyclonal stable lines were maintained in DMEM containing 10% FBS in the presence of 400 μ g geneticin (Gibco).

Mock-transfected or APP695^{LDN}-expressing CHO cells were obtained by stable transfection of pcDNA4 empty vector (Control), wild type hAPP695 cDNA or harboring the London mutation (APP^{LDN}: APPV642I) and subcloned in pcDNA₄ vector. Single clones were maintained in DMEM containing 10% FBS, sodium hypoxanthine-thymidine supplement, and 300 μ M proline as already described [23].

Cells were treated overnight (20 h) with β - or γ -secretase inhibitors. γ -secretase inhibitor ELND006 was used at 5 μ M final concentration and vehicle (methylcellulose/polysorbate 80; Sigma-Aldrich) was used as control [24, 25]. β -secretase inhibitor (Eli Lilly's inhibitor LY288672 [26], synthesized by Elan Pharmaceutical) was used at 30 μ M final concentration prepared in DMSO. Vehicle was used as control since no difference was observed as compared to cells treated with DMSO.

Mice

We used brains isolated from double-transgenic APP₂₃xPS₄₅ mice (B6-Tg *hAPP751 K670N/M671L* \times *PS1 G384A* mThy1). These mice were obtained through Material Transfer Management from Novartis. To generate APP₂₃xPS₄₅ mice, we crossed APP₂₃ transgenic line overexpressing hA β PP with the Swedish double mutation at positions 670/671 [*hAPP751: K670N/M671L*], with PS₄₅ transgenic line overexpressing human PS1 carrying G384A mutation [*hPS1: G384A*] [27]. Both lines are driven by a Thy-1 promoter and were developed in *C57BL/6* genetic background. Control wild type (WT) mice were from the same genetic background.

Ethics statements

In vivo experiments were performed in accordance with relevant guidelines and regulations established by the European community council (Directive of November 24, 1986), and approved by Nice University Animal care and use Committee, and the National Council on animal care of the Ministry of Health (Project n°: NCE/2013-152). Mice were housed in SPF animal facility.

Cellular fractionation

Cell fractionation was performed using SH-SY5Y and CHO cell lines, and brains dissected from WT, APP₂₃, PS₄₅, and APP₂₃PS₄₅ mice using an optimized protocol to isolate subcellular fractions from tissues and cells [28]. Cells ($\approx 10^9$) were harvested, washed by centrifugation at $500 \times g$ for 5 min with PBS. Cells or brains ($n=5$ for each mice group) were resuspended in homogenization buffer (225 mM mannitol, 75 mM sucrose, 30 mM Tris-HCl pH 7.4, 0.1 mM EGTA) and gently disrupted by Dounce homogenization at 4°C. The homogenate was centrifuged twice at $600 \times g$ for 5 min to remove nuclei and unbroken cells, and then the supernatant was centrifuged at $10\,300 \times g$ for 10 min to pellet crude mitochondria (Mc). The resultant supernatant was centrifuged at $20\,000 \times g$ for 30 min at 4°C. The pellet consists of lysosomal and plasma membrane fractions. Further centrifugation of the obtained supernatant at $100\,000 \times g$ for 90 min (70-Ti rotor; Beckman, France) at 4°C results in the isolation of ER (pellet) and cytosolic fraction (supernatant). The crude mitochondrial (Mc) fraction, resuspended in MAM isolation buffer (250 mM mannitol, 5 mM HEPES pH 7.4 and 0.5 mM EGTA), was subjected to Percoll gradient centrifugation (Percoll medium: 225 mM mannitol, 25 mM HEPES pH 7.4, 1 mM EGTA, and 30% vol/vol Percoll) in a 10-ml polycarbonate ultracentrifuge tube. After centrifugation at $95\,000 \times g$ for 30 min (SW40 rotor; Beckman, France), a dense band containing purified mitochondria (Mp) was recovered approximately 3/4 down of the tube, resuspended in MAM isolation buffer and washed by centrifugation at $6300 \times g$ for 10 min to remove the Percoll and finally resuspended in small volume of MAM isolation buffer. The MAMs, containing the structural contacts between mitochondria and ER, collected from the Percoll gradient as a diffuse white band located above the mitochondria, were diluted in MAM isolation buffer and centrifuged at $6300 \times g$ for 10 min; then the supernatant was further centrifuged at $100\,000 \times g$ for 90 min (70-Ti rotor, Beckman) to pellet the MAMs fraction.

SDS-PAGE analysis

To detect A β peptide, protein extracts (40 μ g) were incubated with 70% formic acid (Sigma) and Speed Vac evaporated for 40 min. The pellets were dissolved in 1 M Tris pH 10.8, 25 mM Betaine and diluted in 2x Tris-Tricine loading buffer (125 mM

Tris-HCl pH 8.45, 2% SDS, 20% Glycerol, 0.001% Bromophenol blue, and 5% β -mercaptoethanol). Proteins were resolved by 16.5% Tris-Tricine SDS-PAGE, transferred onto PVDF membranes, and incubated overnight with specific antibodies as specified in legends. All the other proteins were resolved by Tris-Glycine SDS-PAGE following standard procedures.

Antibodies

A β , C99, and full length A β PP were detected using 6E10 antibody (Covance) recognizing 1-16 residues of A β . A β PP C-terminal fragments (A β PP CTF: C99 and C83), were detected using A β PP C-terminal antibody (Sigma Aldrich) recognizing 676–695 residues of A β PP. Other antibodies directed toward the following proteins were as follow: Cytochrome C oxidase subunit II (Cox), and RyR (Thermo scientific); VDAC-1 (Calbiochem); HSP60, Calreticulin, and GRP75 (Santa Cruz); Chapronin 10 (Stressgen, Biotech); Nicastrin, Tom20, α -Tubulin, and β -Actin (Sigma Aldrich), SERCA2b (clone IID8), and Aph1 (Thermo Scientific Pierce Products); BACE-1 and Cyp450 (Abcam); PS1-Nter is a generous gift from Gopal Thinakaran.

Detection of A β by ELISA

The concentrations of A β ₄₀ and A β ₄₂ were measured in subcellular fractions by using the respective ELISA kits (Invitrogen) following the manufacturer's instructions. Briefly, 40 μ g of each fraction was treated with 5 M guanidine, 50 mM HCl solution for 3–4 h at room temperature and stored at –20°C overnight. Supernatant obtained after centrifugation at 16,000 rpm for 20 min at 4°C, were diluted at $>1/10$ in D-PBS-BSA 5% and used for ELISA.

In vitro α -secretase assay

α -secretase activity was monitored as already described [29]. Briefly, 20 μ g of each subcellular fraction were incubated in 90 μ l final volume of Tris buffer (10 mM, pH 7.5) containing α -secretase substrate (JMV2770, 100 μ M) in the absence or presence of the metal chelator *o*-phenanthroline (100 μ M) reported to block metalloproteases. α -secretase activity corresponds to the α -secretase inhibitor-sensitive fluorescence recorded at 320 and 420 nm as excitation and emission wavelengths, respectively.

In vitro β -secretase assay

β -secretase activity was monitored as already described [22]. Briefly, 20 μ g of each subcellular fraction were incubated in a final volume of acetate buffer (25 mM, pH 4.5, 100 μ l) containing BACE-1 substrate [(7-methoxycoumarin-4-yl)-acetyl-SEVNLDAEFRK (2,4-dinitrophenyl)-RRNH₂; 10 μ M; R&D Systems] in the absence or presence of β -secretase inhibitor I (50 μ M; Promo-Cell). BACE-1 activity corresponds to the β -secretase inhibitor I-sensitive fluorescence recorded at 320 and 420 nm as excitation and emission wavelengths, respectively.

In vitro γ -secretase assay

In vitro γ -secretase assay was assessed as already described [30]. Twenty μ g of each subcellular fraction were resuspended in solubilization buffer (150 mM sodium citrate pH 6.4 containing 3-[(3-cholamidypropyl) dimethylammonio]-2-hydroxy-1-propanesulfonate 1% (v/v)) supplemented with protease inhibitor mixture. All steps were performed at 4°C. Solubilized membranes were diluted once with sodium citrate buffer (150 mM pH 6.4), and with reaction buffer (150 mM sodium citrate pH 6.4, 20 mM dithiothreitol, 0.2 mg/ml BSA, 1 mg/ml egg phosphatidyl choline and 50 μ g/mL recombinant C100-FLAG). The resulting reaction mix were then either incubated over constant agitation for 16 h at 37°C or stored at 4°C (negative controls). Samples were then supplemented with 2x Tris-Tricine loading buffer, boiled for 5 min and subjected to 16.5% Tris-Tricine SDS-PAGE.

Immunofluorescence analysis

Cells grown on 25 mm round coverslips were fixed in PFA 4% solution for 10 min at room temperature. Cells were permeabilized with triton 0.5%, and non-specific binding sites were blocked for 1 h with BSA (5%). Cells were then incubated at 4°C overnight with primary antibodies diluted in BSA (5%). After 3 washes, coverslips were incubated with secondary antibodies [fluorescent Alexa Fluor antibodies, Alexa 488- and Alexa 594- or Alexa 633-conjugated (Invitrogen; 1:1000)] at room temperature during 1 h. Immunofluorescence images were acquired on Leica SP5 confocal microscope using excitation filters 488, 594, and 633 nm. Images were analyzed using ImageJ.

High resolution imaging

To visualize mitochondria and ER organelle colocalization, we used GFP construct specifically targeted to the ER. Mitochondria were labeled by using 10 nM Mitotraker[®] Deep Red dye. Z-series images acquired on Leica SP5 microscope were deconvolved. ER and mitochondria colocalization was calculated as the number of voxels (volume pixels) occupied by both signals (i.e., eGFP and Mitotraker Deep Red) over all voxels occupied by the mitochondria (i.e. Mitotraker[®] Deep Red signal) in thresholded images [31]. The spectral proprieties of these two fluorochromes allow specific identification of the two compartments and quantification of contact sites.

Electron microscopy

Cells were fixed *in situ* with 2.5% glutaraldehyde in 0.1 M phosphate buffer at room temperature (RT) for 20 min. Samples were rinsed and then post-fixed with 1% osmium tetroxide and 1% potassium ferrocyanide in 0.1 M cacodylate buffer for 1 h at RT to enhance the staining of cytoplasmic membranes. Cells were rinsed with distilled water, embedded in epoxy resin, sectioned, and examined with a Philips CM12 transmission electron microscope equipped with an Olympus SIS CCD camera. Images were acquired randomly by a blinded investigator. We counted the total number of mitochondria ($n > 150$) in contact or not with ER in more than 10 different micrographs obtained from two different experiments. Data are presented as a percentage of mitochondria in contact with ER.

Neutral lipid staining

Cells grown on 25 mm round coverslips were fixed in PFA 4% solution for 20 min at room temperature. Fixed cells were gently rinsed with PBS 3 times to remove residual formaldehyde. Cells were stained with 1X LipidTOX[™] prepared in PBS for 30 min at room temperature. Coverslips were then processed for image acquisition on Leica SP5 confocal microscope using excitation filters 488. Images were thresholded, converted to mask and analyzed using Particle analysis ImageJ plugin.

Oligomeric A β preparation

Human synthetic A β _{1–42} (Bachem AG, Switzerland) was suspended in hexafluoroisopropanol,

HFIP; Sigma–Aldrich) to 1 mM. Peptide samples were vortexed to obtain a homogenous solution, aliquoted into microfuge tubes, and lyophilized overnight under the hood. The A β _{1–42} peptide films were stored desiccated at –20°C until further processed. To form A β oligomers, peptide films were resuspended to 5 mM in DMSO, diluted in ice cold PBS. Aggregation was allowed to proceed for 24 h at 4 °C before the peptide solution was centrifuged at 14000 \times g for 10 min at 4°C. Supernatant containing oligomeric A β was then used at \approx 5 μ M.

Immunoprecipitation and in-gel digestion

After preclearing with protein A-agarose (Sigma-Aldrich) (1 h, 4°C), MAM fractions (40 μ g) isolated from control CHO cells and overexpressing APP_{LDN} were immunoprecipitated using 6E10 antibody (5 μ g). Immunoprecipitates were recovered by overnight incubation at 4°C, and then incubated with protein A-agarose for 3 h at 4°C. Beads were washed three times with CHRIS buffer (50 mM Tris pH 8, 10% glycerol, 200 mM NaCl, 0.5% Nonidet p-40, and 0.1 mM EDTA) supplemented with protease inhibitors (Complete, Roche diagnostics) and once with PBS. Beads were then resuspended in Laemli 2x buffer, heated at 95°C, and then run on Tris-Glycine PAGE. Gels were stained using Imperial Protein Stain (Thermo scientific), a ready-to-use colorimetric stain formulated with Coomassie dye.

Gel pieces containing proteins were excised and destained by adding 100 μ L of H₂O/ACN solvent (1/1). After 10 min incubation with vortexing the liquid was discarded. This procedure was repeated 2 times. Gel piece were then rinsed (15 min) with acetonitrile and dried under vacuum. Gel pieces were reswelled in 50 μ L of 20 mM dithiothreitol (DTT) in NH₄HCO₃ 100 mM, incubated for 30 min at 56°C, and next cooled down to RT. The DTT solution was replaced with 50 μ L of 55 mM of iodoacetamide in 100 mM of NH₄HCO₃. After 15 min incubation in the dark at room temperature, the solution was discarded and gel pieces were washed by adding successively i) 100 μ L of H₂O/ACN (1/1), repeated 2 times and ii) 100 μ L of acetonitrile. Next gel pieces were reswelled in 50 μ L of 50 mM NH₄HCO₃ buffer containing 10 ng/ μ L of trypsin (modified porcine trypsin sequence grade, Promega) incubated for 1 h at 4°C. Then the solution was removed and replaced by 50 μ L of 50 mM NH₄HCO₃ buffer (without trypsin), and incubated 18 h at 37°C. After trypsin digestion the solution was transferred into an Eppendorf tube

and tryptic peptides were extracted with i) 50 μ L of 1% AF (acid formic) in water (10 min at RT) and ii) 50 μ L acetonitrile (10 min at RT). Peptides extracts were pooled, concentrated under vacuum and solubilized in 15 μ L of 0.1% TFA (trifluoroacetic acid) in water.

NanoHPLC-MALDI-TOF/TOF analysis

Peptide separation was carried out using a nanoHPLC offline (ultimate 3000, Thermo Fisher Scientific) coupled with a MALDI-TOF/TOF mass spectrometer (4800 *plus*, Applied Biosystems). Peptides solution was concentrated on a μ -Precolumn Cartridge Acclaim PepMap 100 C₁₈ (i.d. 5 mm, 5 μ m, 100Å, Thermo Fisher Scientific) at a flow rate of 20 μ L/min and using solvent containing H₂O/ACN/TFA 98%/2%/0.04%. Next peptides separation was performed on a 75 μ m i.d. \times 150 mm (3 μ m, 100Å) Acclaim PepMap 100 C₁₈ column (Thermo Fisher Scientific) at a flow rate of 200 nL/min and with detection at 214 nm. Solvent systems were: (A) 100% water, 0.05% TFA, (B) 100% acetonitrile, 0.04% TFA. The following gradient was used $t=0$ min 100% A; $t=3$ min 100% A; $t=63$ min, 80% B; $t=64$ min, 100% B; $t=69$ min 100% B (temperature was regulated at 30°C).

For offline nanoHPLC-MALDI-TOF/TOF-MS and MS/MS analyses, fractions were collected on an Opti-TOF LC/MALDI target (123 \times 81 mm, Applied Biosystems) and fractionation was done using the Probot fractionation robot (DIONEX, LC Packings). Matrix solution (α -cyano-4-hydroxycinnamic acid, 2.5 mg/mL in 50% water, 50% acetonitrile, 0.1% TFA solution) and nanoHPLC fractions were mixed (in rate 4:1, matrix:fractions) and collected every 20 s (208 fractions were collected per run).

MS spectra were recorded automatically in a mass range of 700–4000 Da resulting from 200 laser shots of constant intensity. Data were collected using 4000 series Explorer (Applied Biosystems) allowing for an automatic selection of peptide masses for subsequent MS/MS experiments. Each MS/MS spectra acquired using 1000 laser shots were further processed using 4000 series Explorer. Finally all raw data were transferred into ProteinPilot software (Applied Biosystems, MDS Analytical Technologies) and proteins identification was processed using ParagonTM Algorithm. Our analyses took in consideration only peptides with confidence index 95%. This experiment was done twice.

Statistical analyses

Results are reported from at least three different experiments or as indicated. Statistical analyses were done using Student's *t*-test, or ANOVA one-way and Tukey post-test or Newman-Keuls Multicomparison post-test. *p* value <0.05 was considered significant.

RESULTS

A β PP and its proteolytic products are present in MAMs fraction of in vitro and in vivo study models of AD

We first aimed at establishing by subcellular fractionation, the distribution of A β PP and its proteolytic catabolites in human SH-SY5Y neuroblastoma cells stably expressing pcDNA3.1 (control) or human A β PP harboring the double Swedish mutations (APP_{swe}: APPKM670/671NL) constructs. We already reported that SH-SY5Y cells expressing APP_{swe} harbor increased A β PP processing yielding the production of A β PP CTFs fragments (C99 and C83) and of A β peptides [22].

We investigated the presence of A β PP and its proteolytic catabolites in MAMs. Although several techniques are available to isolate mitochondria, only few are specifically tuned to the isolation of MAMs, containing unique regions of ER membranes attached to the outer mitochondrial membrane and mitochondria without contamination from other organelles (i.e., pure mitochondria). We used an optimized protocol to isolate these fractions from tissues and cells [28]. Subcellular fractionation procedure was validated by confirming the expression of different proteins in cellular compartments where they were known to be enriched, namely in the ER (SERCA2b, calreticulin (CRT)), in mitochondria (cytochrome oxidase subunit II (Cox), and voltage dependent anion channel 1 (VDAC1)), and in MAMs (Glucose regulated protein 75 (GRP75)) (Fig. 1A, B). Fraction enrichment was further demonstrated by the loss of actin signal in pure mitochondria (Mp), ER, and MAMs fractions (Fig. 1A, B). We observed A β PP and A β expressions in the whole homogenate (homog), in crude mitochondria (Mc), in pure mitochondria (Mp), and in ER fractions (Fig. 1B). Interestingly, we noticed the presence of full-length A β PP and of A β in MAMs fraction of APP_{swe} expressing cells (Fig. 1B). Full-length A β PP was also observed in MAMs fraction of control cells, thus indicating that both endogenous and overex-

pressed A β PP are present in MAMs. However, only a slight A β signal was noticed in control cells. This may be related to low expression level of A β PP in control cells.

We then investigated A β PP subcellular localization in a second cellular model expressing the London APP mutation (APP_{LDN}: APPV642I). In fact, while the Swedish mutation increases the rate of β -cleavage and is proposed to alter the trafficking of A β PP [32], the "London" mutation increases the relative amount of the more toxic A β ₄₂ produced by γ -cleavage, but is not proposed to alter trafficking [33]; Furthermore, It was already reported that CHO cells expressing APP_{LDN} yield high levels of intracellular and secreted A β oligomers [23, 34], and several lines of evidence concord to demonstrate that at least part of the AD-related neurodegenerative process could be due to A β oligomers [34]. Thus, we carried out similar analyses on control and APP_{LDN} cells (Supplementary Figure 1A). We revealed the presence of full length A β PP, and its C-terminal fragments (C99 and C83), and of A β in the whole homogenate, and in Mc, ER, and MAMs fractions of CHO cells expressing APP_{LDN} (Supplementary Figure 1A). Of most interest, we also revealed supplementary bands (Supplementary Figure 1A, see (*) symbol) likely corresponding to dimeric and low molecular oligomeric forms of A β in the Mc and MAMs fractions of CHO cells expressing APP_{LDN} (Supplementary Figure 1A). In this model, we also observed the presence of full-length A β PP in MAMs fraction of control cells. These first set of data show that A β PP, CTFs, and A β are present in MAMs in two *in vitro* models of AD.

We then examined whether A β PP and A β could also be present in MAMs, *in vivo*. We measured expressions of A β PP and its catabolites C99, C83, and A β in subcellular fractions prepared from brains isolated from WT and transgenic mice expressing human A β PP harboring the double Swedish mutations (APP_{swe}: APP₂₃), human presenilin 1 (PS1) (component of the γ -secretase complex) mutation (PS₄₅: G384A), or both (APP₂₃xPS₄₅). Cytochrome p450 (Cyp450, an ER resident protein (Fig. 1A) was indeed enriched in the ER fraction (Fig. 1C). Chapronin 10 (Cpn10), a mitochondrial protein (Fig. 1A) was detected in the homogenate and Mc, and largely enriched in Mp fraction (Fig. 1C). Both APP₂₃ and APP₂₃xPS₄₅ showed A β PP, its C-terminal fragments (C99 and C83), and A β expression in MAMs fraction (Fig. 1C). Interestingly, we also noticed the presence of C83 but to a lesser extent than

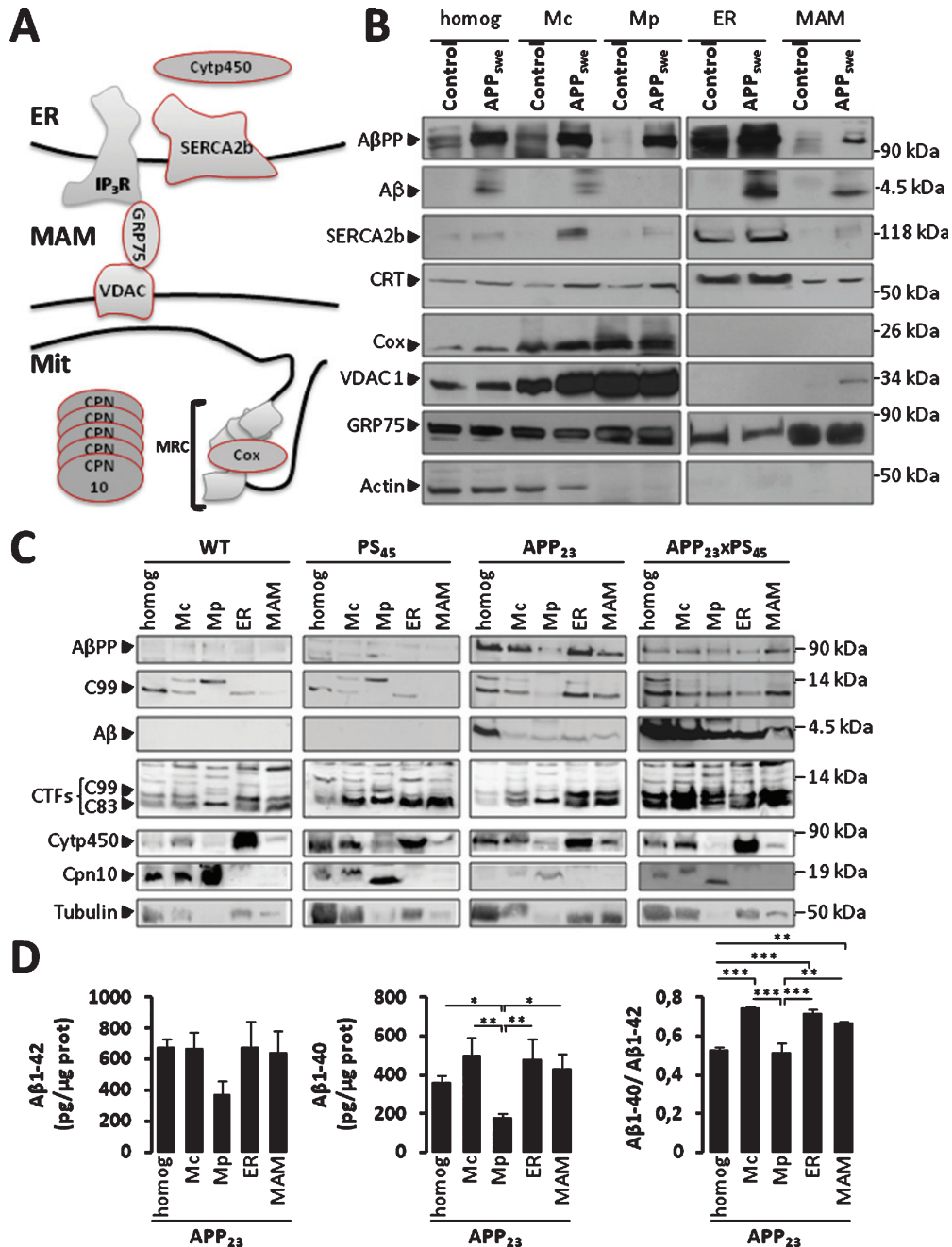


Fig. 1. A β PP and its catabolites are present in MAMs fraction. A) Scheme representing ER-mitochondria contact sites and the molecular markers of ER and mitochondria used. MRC, mitochondrial respiratory chain. B, C) Representative SDS-PAGE of subcellular fractions isolated from: (B) SH-SY5Y neuroblastoma cells expressing pcDNA3.1 empty vector (Control) or hAPP695 harboring double Swedish mutations (APPKM670/671NL: APP^{Swe}), and (C) brains of WT mice, PS₄₅, APP₂₃, and APP₂₃xPS₄₅ transgenic mice. We load 40 μ g (B) or 20 μ g (C) of total homogenate (homog), Crude mitochondria (Mc), pure mitochondria (Mp), the endoplasmic reticulum (ER), and MAMs fractions. A β PP, C99, and A β were detected by 6E10 antibody. A β PP C-terminal fragments (C99 and C83) were detected by A β PP C-ter antibody. SERCA2b, calreticulin (CRT), and Cytochrome p450 (Cyp450) were used as loading controls for ER fraction. Cox, VDAC1, and Chapronin 10 (Cpn10) were used as loading controls for Mc and Mp fractions. GRP75 was used as loading controls for MAMs fraction. Actin and Tubulin were used as loading controls. Data are representative of 3 independent experiments for (B), and were obtained from 5 animals per group of mice and repeated twice for (C). D) ELISA of A β ₄₀ and A β ₄₂ done on subcellular fractions isolated from APP₂₃ transgenic mice. Graph represents Mean \pm S.E.M from three experiments and expressed in pg per μ g of proteins. **p* value <0.05, ***p* value <0.01, ****p* value <0.001 calculated using ANOVA one way and Newman-Keuls Multicomparison post-test.

C99 in WT and PS₄₅ mice (Fig. 1C). The increased C99 fragment in APP₂₃ and APP₂₃PS₄₅ is likely linked to the overexpression of the Swedish mutation known to increase the rate of β -cleavage [32]. Finally specific ELISA approaches allowed unraveling both A β ₁₋₄₀ and A β ₁₋₄₂ peptides in MAMs fraction of APP₂₃ (Fig. 1D). We also unraveled a slightly elevated A β ₄₀/A β ₄₂ ratio in Mc, ER, and MAMs as compared to homogenate and Mp fractions. Overall, this set of data indicates that A β PP and its catabolites are present in MAMs *in vivo* (Fig. 1C, D) in agreement with the *in vitro* data (Fig. 1B and Supplementary Figure 1).

Immunolocalization of A β PP and its catabolites in the ER, mitochondria, and MAMs

We then analyzed subcellular localization of A β PP and its catabolites by immunofluorescence in SH-SY5Y cells expressing APP_{swe}. We investigated the extent of A β PP, and its catabolites co-localization with the ER and mitochondria by using A β PP C-ter antibody (recognizing full length A β PP, C-terminal fragments (C83 and C99), and A β PP intracellular domain (AICD)), or 6E10 antibody (recognizing full length A β PP, C99, and A β peptides) concomitantly with specific antibodies recognizing proteins localized in the ER (the ryanodine receptor (RyR) and Cyp450), or in the mitochondria (Tom20 and Cox). Representative images show the co-localization (yellow signal) of A β PP and its catabolites with both the ER (Fig. 2A, C), and mitochondria (Fig. 2B, D) markers. Similar results were obtained in CHO cells expressing APP_{LDN} using A β PP C-ter antibody (Supplementary Figure 1B, C).

We then examined A β PP and its catabolites colocalization within ER-mitochondria contact sites. SH-SY5Y cells expressing APP_{swe} were immunolabeled concomitantly with A β PP C-ter, Tom20, and the ER-resident protein calreticulin (CRT) antibodies (Fig. 2E). Merge, high magnification images (Fig. 2E, merge and inset), and super imposable line scan traces (Fig. 2F) (corresponding to the red, green, and blue fluorescence signals along the red line in the inset) demonstrate co-localization of A β PP and CTFs with both ER and mitochondria. These data are in agreement with our data obtained by subcellular fractionation and further support the presence of A β PP and its metabolites in MAMs in cellular models overexpressing both the Swedish and the London mutations.

Full length A β PP as revealed by using specific antibody recognizing A β PP N-terminal epitope (APP N-ter) shows also slight co-localization with the ER (Supplementary Figure 3A) and mitochondria (Supplementary Figure 3B).

Using immunofluorescence and imaging, we also examined the subcellular distribution of wild type A β PP in both SH-SY5Y and CHO cells. We show the immunolocalization of WT A β PP and its metabolites in the ER (colocalization with SERCA2b) (Supplementary Figures 3A and 4A), mitochondria (colocalization with mitochondrial HSP60) (Supplementary Figures 3B and 4B), and MAMs (Supplementary Figures 3C, D and 4C, D) in both SH-SY5Y (Supplementary Figure 3) and CHO (Supplementary Figure 4) cells overexpressing wild type A β PP. The results reveal that like the Swedish and London mutated A β PP, wild type A β PP harbor partial colocalization with the ER, mitochondria, and MAMs. These data strengthen our conclusion regarding the subcellular localization and proteolytic processing of A β PP in MAMs.

*β - and γ -secretases are present and show *in vitro* activities in MAMs fraction*

We next revealed by subcellular fractionation of APP₂₃ transgenic mice brains the presence of β -secretase (BACE-1) and of components of the γ -secretase complex (PS1, Aph1, and nicastrin) in MAMs (Fig. 3A). *In vitro* γ -secretase activity assay allowed us to reveal the hydrolysis of recombinant C100 fragment at 37°C and not at 4°C (used as negative control) that yield A β production in the homogenate, Mc, Mp, ER, and MAMs fractions isolated from brains of APP₂₃ mice (Fig. 3B). We did not notice a difference in γ -secretase activity between WT and APP₂₃ mice (Supplementary Figure 5A). Similarly, γ -secretase activity was also observed in MAMs fraction isolated from CHO cells expressing APP_{LDN} (Supplementary Figure 5B). In agreement with BACE-1 expression in APP₂₃ MAMs (Fig. 3A), we detect β -secretase activity in MAMs fraction in both WT and APP₂₃ mice fractions (Fig. 3C), and in control CHO cells and expressing APP_{LDN} (Supplementary Figure 5C). Related to the presence of C83 (the α -secretase-derived CTF) *in vivo* (Fig. 1C), we also measured α -secretase activity in subcellular fractions isolated from APP₂₃ transgenic mice. We indeed could observe *in vitro* α -secretase activity in the homogenate, Mc, ER, and MAMs. However, the *in vitro* α -secretase activity was lower in MAMs

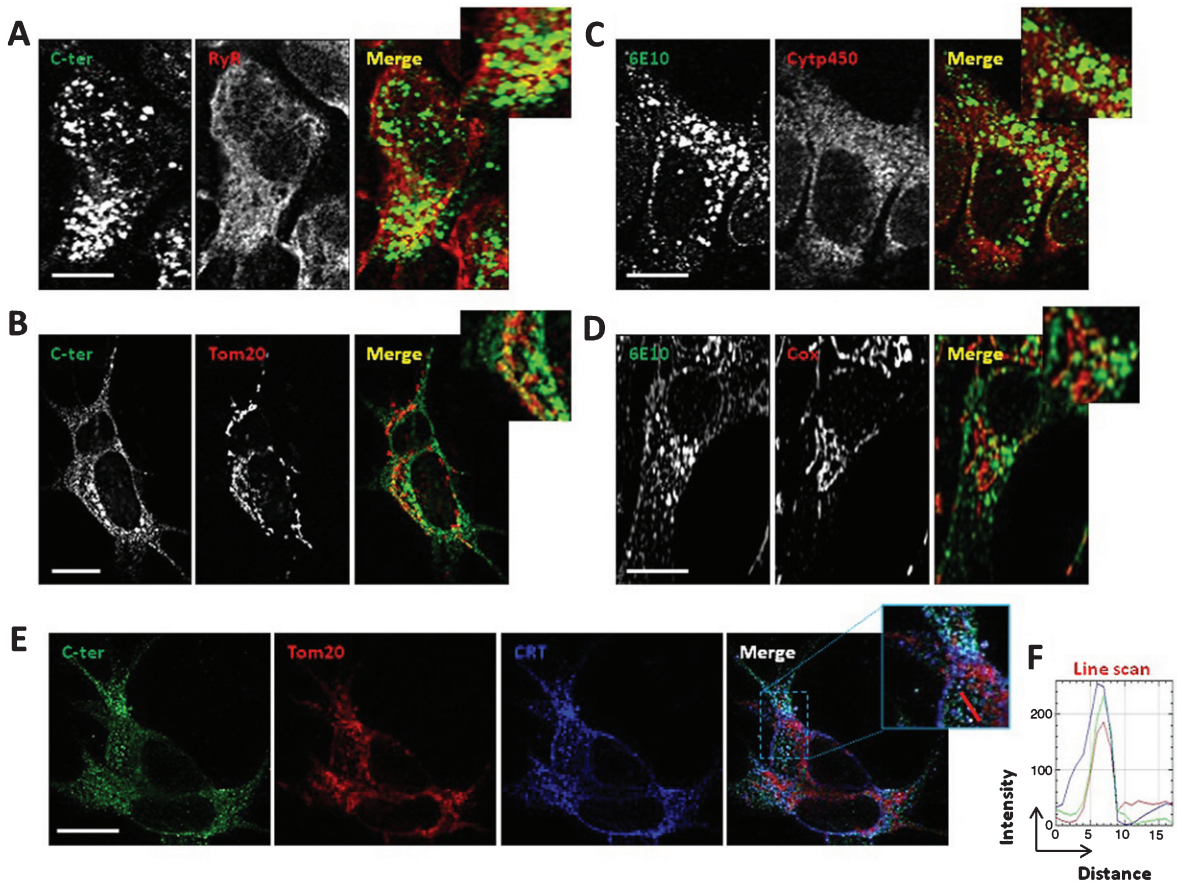


Fig. 2. A β PP and its catabolites colocalize with ER and mitochondria. A, B) Immunostaining of SH-SY5Y APP_{swe} cells with A β PP C-ter antibody recognizing total A β PP and CTF (C99 and C83). RyR was used to stain the ER (A). Tom20 was used to stain mitochondria (B). C, D) Immunostaining of SH-SY5Y APP_{swe} cells with 6E10 antibody recognizing total A β PP, C99, and A β peptides. Cytp450 was used to stain the ER (C). Cox was used to stain mitochondria (D). A-D) Merge images show green and red signals and were used to show the colocalization of A β PP and its catabolites with the ER or mitochondria (yellow signal). Scale bars represent 10 μ m. E) Immunostaining of SH-SY5Y APP_{swe} cells with A β PP C-ter, Tom20, and calreticulin (CRT) antibodies. Merge images show overlay of green, red and blue signals, where colocalization is depicted in white (see merge image and magnified overlay). Scale bar represents 10 μ m. F) RGB (Red, Green, Blue) profile plots of a representative line scan (red line in inset) shows the maximum intensity of Red, Green, and Blue plots observed at the same distance, and demonstrating the colocalization of Red (Tom20), Green (C-ter), and Blue (CRT) signals.

as compared to the other fractions (Supplementary Figure 5D). All over, and according to already published data [9, 13, 16, 20], we revealed the presence and activity of γ -secretase in MAMs fraction. We also provide for the first time evidences showing that BACE-1 is present in MAMs fraction and that BACE-1 and α -secretase may harbor enzymatic activities in this domain.

Increased ER and mitochondria contacts in SH-SY5Y neuroblastoma APP_{swe} cell model

Increased ER-mitochondria colocalization has been reported in human-derived fibroblasts isolated from familial and sporadic AD patients and in mouse

embryonic fibroblasts isolated from either PS1 or PS1 and PS2 knockout mice [21]. This was also observed in cells overexpressing familial AD PS2 mutant [35]. We examined herein ER-mitochondria contact sites in live cells overexpressing APP_{swe} (Fig. 4A). For live cells imaging, the ER was visualized by transfecting GFP construct targeted to the ER (ER GFP). Mitochondria were labeled by using Mitotracker Deep red dye as already described [31]. We noticed that APP_{swe} expressing cells exhibit a significant increased colocalization of the ER and mitochondria as compared to control cells (evidenced by an increase of mitochondria volume in contact with the ER versus total mitochondria volume) (Fig. 4A). Accordingly, we also revealed increased mitochondria in contact with

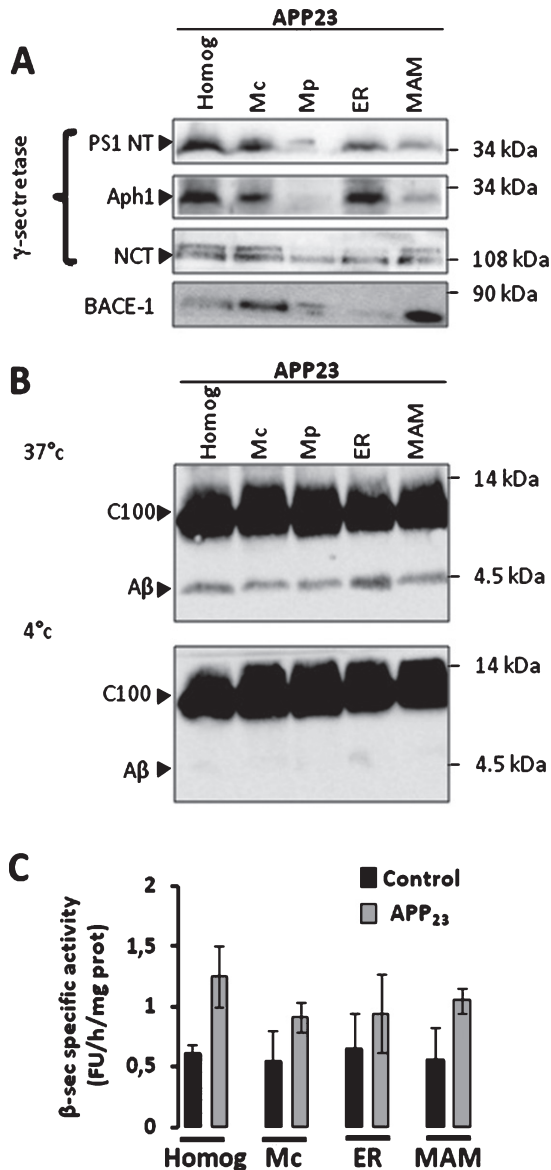


Fig. 3. β -secretase and γ -secretase complex are present and active in MAMs fraction in mice brains. A) Representative SDS-PAGE of subcellular fractions isolated from APP₂₃ transgenic mice as in Fig. 1 showing the expression of γ -secretase complex (cleaved Presenilin 1 N-terminal fragment (PS1-NT); Anterior Pharynx-Defective 1 (Aph1); and Nicastrin (NCT)) and of β -secretase (BACE-1). B) Cell-free A β production from recombinant C100 peptide performed at 37°C or 4°C in the presence of subcellular fractions isolated from APP₂₃ transgenic mice. C100 and A β were detected using 6E10 antibody. C) *In vitro* β -secretase specific activity in WT and APP₂₃ mice brain fractions. The graph represents β -secretase specific activity obtained by subtracting residual activity in the presence of β -secretase inhibitor from total β -secretase activity and is expressed in fluorescent units (FU)/hour (h)/mg of proteins of each fraction. Data are presented as Mean \pm S.E.M. obtained from 3 independent experiments. Differences are statistically non-significant using ANOVA one-way and Newman-Keuls Multicomparison post-test.

the ER by electron microscopy in cells overexpressing APP_{swe} as compared to controls (Fig. 4B).

These data demonstrate that A β PP mutation/overexpression and most likely its proteolytic products lead to increased ER-mitochondria contact sites.

The increased ER-mitochondria contacts in SH-SY5Y neuroblastoma APP_{swe} cell model may depend on mitochondrial or ER shape and volume. Supplementary experiments are necessary to fully study this issue.

Overexpression of A β PP familial mutations increases neutral lipids accumulation

MAM have the proprieties of an intracellular lipid raft [21]. This micro-domain is also the site of phospholipid synthesis and transport between the ER and mitochondria [36]. We explored the potential lipid dysfunction in relation with A β PP and its catabolites presence in the MAM and increased ER and mitochondria contacts. In particular, lipid droplets are found in close proximity with the ER and mitochondria [37, 38].

We used LipidTox GreenTM dye to stain and quantify neutral lipid accumulation (major components of lipid droplets). Whereas the lipidTox stain in control CHO cells reveal few positive droplets, the CHO cells expressing APP_{LDN} contained numerous lipidTox positive droplets (\sim 2-fold over control cells) (Fig. 5A, B). Similar results were also observed in SH-SY5Y neuroblastoma cells expressing APP_{swe} as compared to its respective control cells (Supplementary Figure 6A, B). Importantly, the increase in lipid droplets in CHO cells expressing APP_{LDN} was reversed by inhibition of β - or γ -secretases. As expected, both inhibitors block A β production (Supplementary Figure 6C). However, while the inhibitor of β -secretase strongly reduces C99 fragment production, inhibitor of γ -secretase enhances the accumulation of this fragment (Supplementary Figure 6C). Based on these results, we may postulate that lipid droplets accumulation in our study models is likely linked to the production of A β peptides instead of CTFs.

We further demonstrate the specificity of our data regarding the reduction of neutral lipid accumulation obtained with β - and γ -secretase inhibitors observed in CHO cells expressing the London A β PP mutant. First, we analyzed neutral lipid accumulation in naïve SH-SY5Y cells upon the application of exogenous oligomeric A β (Fig. 5C, 5D). We could indeed

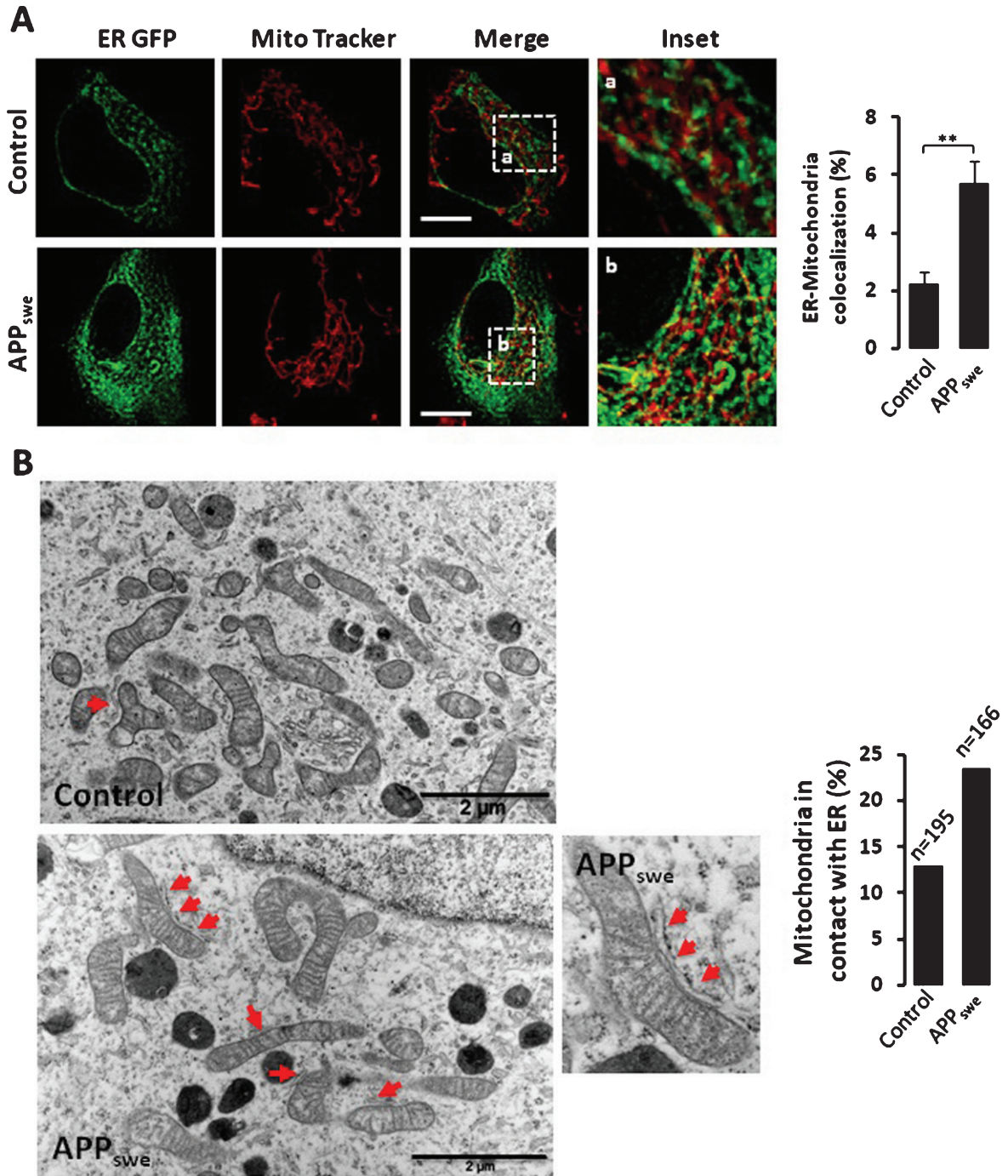


Fig. 4. Increased ER-mitochondria contact sites in SH-SY5Y cells expressing APP_{swe}. A) Quantitative analyses of the colocalization of ER and mitochondria in SH-SY5Y live cells expressing pcDNA3.1 [Control ($n=4$) or APP_{swe} ($n=7$)]. ER is visualized by transfecting ER GFP (green). Mitochondria are stained with Mitotracker Deep red dye (red). Merge images show overlay of green and red signals. Insets (magnified overlay) showing ER and mitochondria colocalization depicted in yellow. The graph represents the quantification of ER and mitochondria colocalization presented as percentage of total mitochondrial volume (% mean \pm S.E.M.). ** p value < 0.01 using Student's t test. Scale bars represent 5 μ m. B) Representative electron microscopy micrographs of control and APP_{swe} expressing cells. Red arrows show mitochondria in contacts with ER. High magnification of ER-mitochondria contacts is shown for APP_{swe} cells. Data are presented as % of mitochondria in contacts with ER. Quantification was obtained from a total number of mitochondria in control ($n=195$) and in APP_{swe} expressing cells ($n=166$). Scale bars represent 2 μ m.

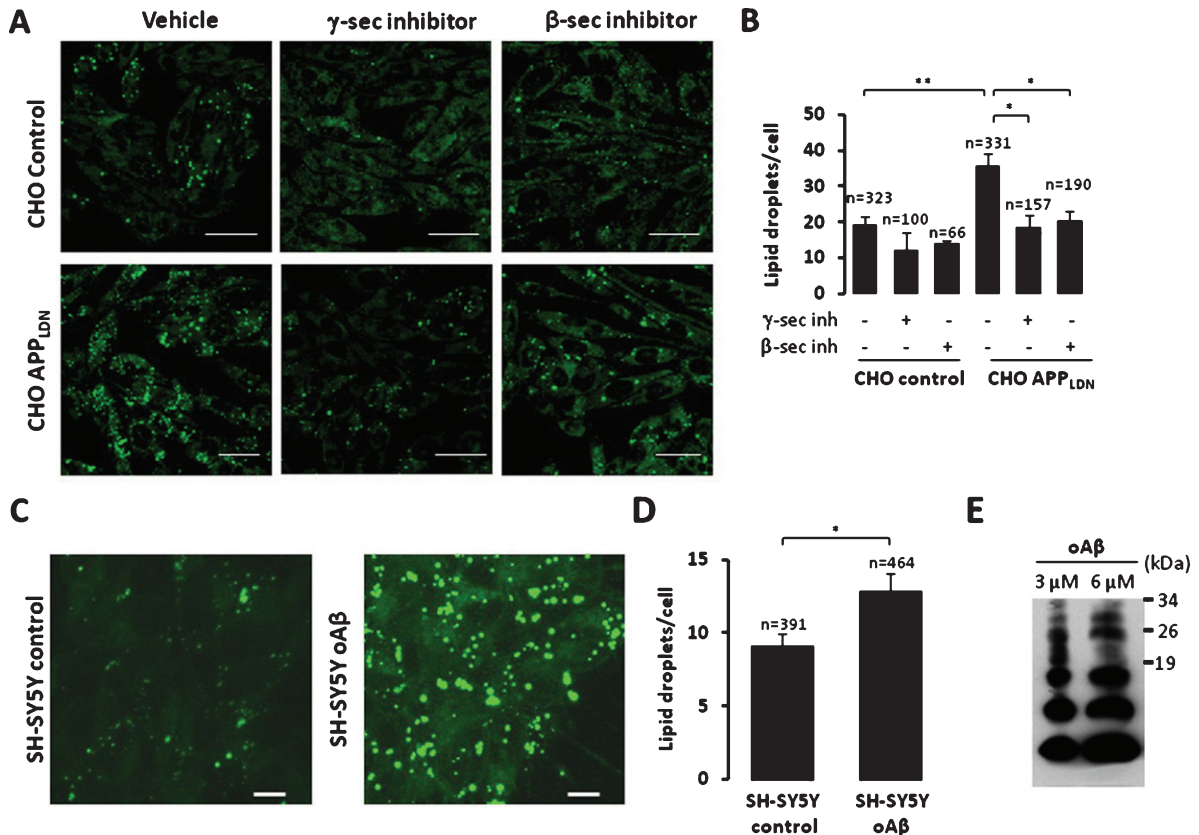


Fig. 5. Increased neutral lipid droplets accumulation in CHO cells expressing APP_{LDN}. A) Representative images showing neutral lipids staining in CHO control cells or expressing APP_{LDN}. Cells were treated with vehicle or with β- or γ-secretase inhibitors for 20 h. Scale bars represent 25 μm. B) The graph shows the quantification of lipid droplets (mean ± S.E.M. number of puncta/cell). The number of analyzed cells is indicated for each condition and was obtained from at least three independent experiments. **p* value < 0.05, and ***p* value < 0.01 using ANOVA one way and Tukey post-test. C) Representative images showing neutral lipids staining in SH-SY5Y cells non treated or treated with 5 μM oligomeric Aβ (oAβ) preparation for 20 h. Scale bars represent 10 μm. D) The graph shows the quantification of lipid droplets (mean ± S.E.M. number of puncta/cell). The number of analyzed cells is indicated for each condition and was obtained from two independent experiments. **p* value < 0.05, using Student *t*-test. E) Representative SDS-PAGE of oligomeric Aβ (3 μM and 6 μM) preparation.

show that the treatment with exogenous oligomeric Aβ leads to increased neutral lipid accumulation (Fig. 5C, D). However, since our Aβ preparation contains both monomeric and oligomeric forms of Aβ (Fig. 5E), we cannot conclude whether neutral lipids accumulation is linked to either monomeric or oligomeric Aβ (Fig. 5E). Second, we analyzed neural lipid accumulation in mouse embryonic fibroblasts (MEF) isolated from wild type (expressing endogenous AβPP) (WT MEF), or invalidated for AβPP and its family member APLP2 (referred to as APP-TKO MEF) [39]. We established that knock out of AβPP did not affect neutral lipid accumulation (Supplementary Figure 6D). Importantly, we did not notice an effect of both β- and γ-secretase inhibitors neither in WT MEF, nor in APP-TKO MEF cells (Supplementary Figure 6D).

All over, these data demonstrates that neutral lipid accumulation is likely linked to AβPP overexpression and processing and to Aβ production.

Protein interactome of AβPP and its catabolites in MAMs

Since we showed that AβPP is present and is processed in MAMs likely contributing to increased ER-mitochondria contact sites, we hypothesized that AβPP and its catabolites may thus interact with proteins localized in this micro-domain. We used immunoprecipitation combined to proteomic approach to reveal AβPP and its catabolites protein interactome in MAMs. By using 6E10 antibody, we expected to immunoprecipitate MAMs proteins interacting directly or indirectly with AβPP holoprotein,

C99, and/or A β peptides. To exclude false positives due to adsorption of MAMs proteins with IgG beads, we incubated MAMs fractions isolated from CHO cells expressing pcDNA4 (Control) or APP_{LDN} with IgG beads alone (Supplementary Figure 7). We validated our experimental procedure by revealing A β PP peptides (accession number: P12023) in MAMs fraction isolated from APP_{LDN} CHO cells but not in CHO control cells, thus providing a positive control of our experimental protocol (Supplementary Figure 8).

We specifically immunoprecipitated 135 proteins from CHO APP_{LDN} MAM fraction (obtained in 6E10 immunoprecipitate conditions, but not in samples incubated with IgG beads alone) (Supplementary Figure 7). In the analyses, we excluded IgG and ribosomal proteins. Among immunoprecipitated proteins from CHO APP_{LDN} MAMs fraction, 39 were also identified in CHO control MAMs fraction. The common proteins identified in both samples may reflect MAMs proteins interacting with full length A β PP rather than those interacting with C99 and A β peptides. Indeed, the amount of A β PP catabolites was clearly lower in control as compared to APP_{LDN} MAMs fraction (Supplementary Figure 1).

We analyzed the subcellular localization and biological functions of the 96 unique proteins obtained in CHO APP_{LDN} MAMs fraction by using gene ontology (GO) annotation and UniProt (Universal Protein Resource) database. In this study, we exclude from the analysis proteins showing nuclear, extracellular or microtubule localization. We then determined that 38 proteins are resident in mitochondria (44%); 23 proteins have ER localization (27%), 12 proteins are mainly cytosolic (14%), 7 proteins are found in lysosomes, Golgi apparatus, and endosomes (8%), and 6 proteins are associated with the plasma membrane (7%) (Supplementary Figure 7). It is important to note that mitochondria and ER comprise 71% of the total immunoprecipitated proteins. However, proteins associated with other organelles should not be discounted as contaminants since the plasma membrane, Golgi apparatus, and lysosomes all have tight association with the ER or mitochondria and may execute fundamental cellular functions in these contact microdomains [40, 41].

We then analyzed the types of function of the proteins identified. Of most interest, the identified molecular functions (Supplementary Figure 7) are mostly attributable to MAMs functions [41, 42]. We focused on the four most significant biological processes: 1) mitochondrial function and transport ($n=23$) (Supplementary Figure 7 and

Table 1); 2) protein synthesis, maturation, folding, processing, and degradation ($n=18$) (Supplementary Figure 7 and Table 2); 3) vesicle formation and trafficking ($n=12$) (Supplementary Figure 7 and Table 3); and 4) fatty-acid metabolism ($n=10$) (Supplementary Figure 7 and Table 4). The remaining pathways correspond to proteins implicated in mitochondria and ER structure ($n=3$), apoptosis ($n=2$), cellular signaling ($n=7$), and other metabolic pathways ($n=4$) (Supplementary Figure 7 and Table 1).

Protein interactome related to mitochondrial function

The identified mitochondrial proteins ($n=23$) are implicated in mitochondrial respiratory chain complexes ($n=9$), mitochondrial transport ($n=6$), and mitochondrial metabolism ($n=8$) (Supplementary Figure 7 and Table 1). Interestingly, 6 out of 10 isolated mitochondrial respiratory chain proteins are components or chaperone proteins of the mitochondrial complex I. We also identified Cytochrome b-c1 complex subunit1 (component of the mitochondrial respiratory chain complex III), and two components of the mitochondrial ATP synthase complex (Supplementary Figure 7 and Table 1). The protein interactome of A β PP and its catabolites related to mitochondrial function include also proteins implicated in mitochondrial metabolism (e.g., Acyl-CoA, and coenzyme Q biosynthesis) (Table 1). In accordance with already reported data, our approach allowed us to identify proteins implicated in mitochondrial transport (e.g., TIM, and TOM complexes) [8, 43, 44] (Supplementary Figure 7 and Table 1). This result could be considered as a positive control validating our experimental procedure.

Protein interactome related to protein synthesis, maturation, folding, processing, and degradation

We identified different proteins implicated in ER protein maturation (e.g., glycosylation and signal peptidase complex), protein folding (e.g., inhibition of protein aggregation of misfolded proteins, protein quality control), and protein processing and degradation (e.g., peptidases, proteases) (Supplementary Figure 7 and Table 2). The protein interactome includes also chaperone proteins inhibiting the aggregation of misfolded proteins (e.g., the protein disulfide-isomerase A4, the protein

Table 1
Classification of protein interactome of the A β PP and its catabolites related to mitochondrial function

UniProt ID	Name ^(a)	Peptides (95%) ^(b)	Function
<i>Mitochondrial respiratory chain complexes</i>			
Q9CZ13	Cytochrome b-c1 complex subunit 1, mitochondrial GN = Uqcrc1	3	Component of complex III or cytochrome b-c1 complex
Q8BGH2 ^(*)	Sorting and assembly machinery component 50 homolog GN = Samm50	2	Maintenance of the structure of mitochondrial cristae and the proper assembly of the MRC complexes
Q91VD9	NADH-ubiquinone oxidoreductase 75 kDa subunit, mitochondrial GN = Ndufs1	1	Core subunit of the mitochondrial Complex I
Q91WD5	NADH dehydrogenase [ubiquinone] iron-sulfur protein 2, mitochondrial GN = Ndufs2	1	Core subunit of the mitochondrial Complex I
Q91YT0	NADH dehydrogenase [ubiquinone] flavoprotein 1, mitochondrial GN = Ndufv1	1	Core subunit of the mitochondrial Complex I
Q9CQN1	Heat shock protein 75 kDa, mitochondrial GN = Trap1	1	Chaperone protein involved in maintaining mitochondrial function and polarization, likely through stabilization of mitochondrial complex I
Q9CWX2	Complex I intermediate-associated protein 30, mitochondrial GN = Ndufaf1	1	Chaperone protein involved in the assembly of the mitochondrial complex I
P97450	ATP synthase-coupling factor 6, mitochondrial GN = Atp5j	1	Mitochondrial F ₁ F ₀ ATP synthase or Complex V
Q9CQQ7	ATP synthase subunit b, mitochondrial GN = Atp5f1	1	Mitochondrial F ₁ F ₀ ATP synthase or Complex V
<i>Mitochondrial transport</i>			
Q9Z0V7	Mitochondrial import inner membrane translocase subunit Tim17-B GN = Timm17b	1	Essential component of the TIM23 complex
Q8BGH2 ^(*)	Sorting and assembly machinery component 50 homolog GN = Samm50	2	Assembly of TOM40 into the TOM complex
Q9CPQ3	Mitochondrial import receptor subunit TOM22 homolog GN = Tomm22	1	Central receptor component of the translocase of the TOM complex
Q9QZD8	Mitochondrial dicarboxylate carrier GN = Slc25a10	1	Translocation of malonate, malate and succinate in exchange for phosphate, sulfate, sulfite or thiosulfate
Q9Z2Z6	Mitochondrial carnitine/acylcarnitine carrier protein OS = Mus musculus GN = Slc25a20	1	Transport of acylcarnitines of different length across the mitochondrial inner membrane
Q9CQX2	Cytochrome b5 type B GN = Cyb5b	1	Electron carrier for several membrane bound oxygenases
<i>Mitochondrial metabolism</i>			
Q9JHI5	Isovaleryl-CoA dehydrogenase, mitochondrial GN = Ivd	1	Synthesis of (S)-3-hydroxy-3-methylglutaryl-CoA from 3-isovaleryl-CoA
Q9WUM5	Succinyl-CoA ligase [ADP/GDP-forming] subunit alpha, mitochondrial GN = Suc1g1	1	ATP- or GTP-dependent ligation of succinate and CoA to form succinyl-CoA
Q9R112	Sulfide:quinone oxidoreductase, mitochondrial GN = Sqrd1	1	Oxidation of hydrogen sulfide with the help of a quinone, such as ubiquinone
Q9D2G2	Dihydrolipoyllysine-residue succinyltransferase component of 2-oxoglutarate dehydrogenase complex, mitochondrial GN = Dlst	1	Conversion of 2-oxoglutarate to succinyl-CoA and CO ₂
Q60936	Chaperone activity of bc1 complex-like, mitochondrial GN = Adck3	1	Biosynthesis of coenzyme Q (ubiquinone)
Q64521	Glycerol-3-phosphate dehydrogenase, mitochondrial GN = Gpd2	1	Belongs to the FAD-dependent glycerol-3-phosphate dehydrogenase family
Q99KI0	Aconitate hydratase, mitochondrial GN = Aco2	1	Isomerization of citrate to isocitrate via cis-aconitate
Q9D051	Pyruvate dehydrogenase E1 component subunit beta, mitochondrial GN = Pdhh	1	Conversion of pyruvate to acetyl-CoA and CO ₂

UniProtID, protein accession number in Swiss-Prot/Trembl. ^(a)Protein name given by UniProt database. ^(b)Number of peptides matching the protein with a confidence limit of 95%. ^(*)A protein that have several functions. GN, gene name.

Table 2

Classification of protein interactome of the A β PP and its catabolites related to protein synthesis, maturation, folding, processing and degradation

UniProt ID	Name ^(a)	Peptides (95%) ^(b)	Function
<i>Protein synthesis</i>			
Q8K0D5	Elongation factor G, mitochondrial GN = Gfm1	2	Mitochondrial GTPase that catalyzes the GTP-dependent ribosomal translocation step during translation elongation.
Q8R2Y8	Peptidyl-tRNA hydrolase 2, mitochondrial GN = Ptrh2	2	The natural substrate for this enzyme may be peptidyl-tRNAs which drop off the ribosome during protein synthesis.
<i>Protein maturation or modification</i>			
Q9DBG6	Dolichyl-diphosphooligosaccharide–protein glycosyltransferase subunit 2 GN = Rpn2	1	Involved in the pathway of protein glycosylation
P46978	Dolichyl-diphosphooligosaccharide–protein glycosyltransferase subunit STT3A GN = Stt3a	1	Involved in the pathway of protein glycosylation
Q9R0P6	Signal peptidase complex catalytic subunit SEC11A GN = Sec11a	1	Component of the microsomal signal peptidase complex.
Q9CXY9	GPI-anchor transamidase GN = Pigk	1	Involved in the pathway glycosylphosphatidylinositol-anchor biosynthesis
<i>Protein folding</i>			
P08003	Protein disulfide-isomerase A4 GN = Pdia4	2	Chaperone that inhibits aggregation of misfolded proteins.
Q922R8	Protein disulfide-isomerase A6 GN = Pdia6	2	Chaperone that inhibits aggregation of misfolded proteins.
Q9D1M7	Peptidyl-prolyl cis-trans isomerase FKBP11 GN = Fkbp11	2	PPIases accelerate the folding of proteins during protein synthesis.
Q9Z2G6	Protein sel-1 homolog 1 GN = Sel11	2	May play a role in Notch signaling. May be involved in the ER quality control
P24369	Peptidyl-prolyl cis-trans isomerase B GN = Ppiib	1	PPIases accelerate the folding of proteins
Q99KV1	DnaJ homolog subfamily B member 11 GN = Dnajb11	1	Co-chaperone for HSPA5. Binds directly to both unfolded proteins that are substrates for ERAD
<i>Protein processing and degradation</i>			
Q8K411	Presequence protease, mitochondrial GN = Pitrm1	1	ATP-independent protease that degrades mitochondrial transit peptides after their cleavage
Q6I6G8	E3 ubiquitin-protein ligase HECW2 GN = Hecw2	1	Involved in the pathway of protein ubiquitination,
O70481	E3 ubiquitin-protein ligase UBR1 GN = Ubr1	1	E3 ubiquitin-protein ligase which is a component of the N-end rule pathway
O88696	Putative ATP-dependent Clp protease proteolytic subunit, mitochondrial GN = Clpp	1	Protease component of the Clp complex that cleaves peptides and various proteins in an ATP-dependent process
Q9QXV0	ProSAAS GN = Pcsk1n	1	Serine-type endopeptidase inhibitor activity
Q9CYN2	Signal peptidase complex subunit 2 GN = Spcs2	1	Component of the microsomal signal peptidase complex

UniProtID, protein accession number in Swiss-Prot/Trembl. ^(a)Protein name given by UniProt database. ^(b)Number of unique peptides matching the protein with a confidence limit of 95%. GN, gene name.

disulfide-isomerase A6, and the peptidyl-prolyl cis-isomerase FKBP11), or implicated in the ER quality control (e.g., the protein sel-1 homolog 1, and the DnaJ homolog subfamily B member 11). Interestingly, we also identified proteins implicated in mitochondrial protein translation, and the mitochondrial presequence protease involved in the degradation of mitochondrial transit peptides (Supplementary Figure 7 and Table 2).

Protein interactome related to vesicle formation and trafficking

In addition to the mitochondria, the ER is associated with different subcellular domains that execute divers function such as vesicular traffic [40]. As a matter of fact, we identified a class of A β PP protein interactome ($n=12$) including proteins implicated in vesicle formation and trafficking (Supplementary

Table 3
Classification of protein interactome of the A β PP and its catabolites related to vesicle formation and trafficking

UniProt ID	Name ^(a)	Peptides (95%) ^(b)	Function
Q3V009	Transmembrane emp24 domain-containing protein 1 GN = Tmed1	1	Potential role in vesicular protein trafficking, mainly in the early secretory pathway
Q9R0Q3	Transmembrane emp24 domain-containing protein 2 GN = Tmed2	2	Involved in vesicular protein trafficking (early secretory pathway and post-Golgi membranes)
Q9D1D4	Transmembrane emp24 domain-containing protein 10 GN = Tmed10	1	Involved in vesicular protein trafficking
O08917	Flotillin-1 GN = Flot1	1	May act as a scaffolding protein within caveolar membranes, functionally participating in formation of caveolae or caveolae-like vesicles
O55100	Synaptogyrin-1 GN = Syngr1	1	Involved in the regulation of short-term and long-term synaptic plasticity
P35293	Ras-related protein Rab-18 GN = Rab18	1	Plays a role in apical endocytosis/recycling
P61027	Ras-related protein Rab-10 GN = Rab10	2	Regulator of intracellular membrane trafficking, may play a role in ER dynamics and morphology
Q9D1G1	Ras-related protein Rab-1B GN = Rab1b	1	Regulates vesicular transport between the ER and Golgi compartments. Plays a role in the initial events of the autophagic vacuole development
Q61548	Clathrin coat assembly protein AP180 GN = Snap91	1	Components of the adaptor complexes which link clathrin to receptors in coated vesicles
Q9CQW2	ADP-ribosylation factor-like protein 8B GN = Arl8b	1	May play a role in lysosomes motility
Q9EQH3	Vacuolar protein sorting-associated protein 35 GN = Vps35	1	Component of the retromer cargo-selective complex (CSC).
Q9JIG8	PRA1 family protein 2 GN = Praf2	1	May be involved in ER/Golgi transport and vesicular traffic

UniProtID, protein accession number in Swiss-Prot/Trembl. ^(a)Protein name given by UniProt database. ^(b)Number of unique peptides matching the protein with a confidence limit of 95%. GN, gene name.

Figure 7 and Table 3). These include Ras-related family proteins (Rab-10, Rab18, and Rab1b) implicated in ER-Golgi transport and vesicular traffic, and Praf2 (PRAF1 family protein 2) known to interacts with numerous Rab GTPases thus assisting in the packaging of Rab proteins into vesicles for the transport to target compartments [45, 46]. We also revealed other proteins implicated in vesicular transport namely the transmembrane emp24 domain containing proteins (Tmed2, Tmed10, and Tmed1) and Arl8b (an Arf-like GTPase being a critical regulator of cargo delivery to lysosomes). We also revealed in the protein interactome Flotillin 1, known to be a scaffolding protein within caveolar membranes and to interact with lipid rafts [47, 48].

Protein interactome related to fatty-acid metabolism

We classified a fourth cellular pathway including proteins ($n = 10$) implicated in fatty-acid metabolism (Supplementary Figure 7 and Table 4). This includes proteins implicated in fatty-acid elongation or

modification (Hsd17b12, Acot1, Far1), fatty-acid beta-oxidation (Acaa2, Hadhb, and Acad9), sterol and cholesterol biosynthesis (Nsdh1, Cyp51a1, Lss), and sphingolipid metabolism (Sptlc1). Based on GO annotation, these proteins are mostly localized in the ER (Hsd17b12, Nsdh1, Sptlc1, Cyp51a1, and Lss), or in the mitochondria (Acot1, Acaa2, Far1, and Hadhb, and Acad9). Nsdh1 may be also localized in lipid droplets (Table 4).

All over, these analyses identify potential new A β PP and its catabolites interactors in MAMs. However, a dedicated study is necessary to confirm specific direct or indirect interactions of A β PP or its catabolites with these proteins and their role in MAMs alteration in AD.

DISCUSSION

We used both biochemical and imaging approaches to demonstrate the presence of A β PP and its catabolites in MAMs in *in vitro* and in *in vivo* AD-like models. It is well established that A β PP is synthe-

Table 4
Classification of protein intercatome of the A β PP and its catabolites related to fatty-acid metabolism

UniProt ID	Name ^(a)	Peptides (95%) ^(b)	Function
O70503	Estradiol 17-beta-dehydrogenase 12 GN = Hsd17b12	2	Catalyzes the second of the four reactions of the long-chain fatty acids elongation cycle
Q9R1J0	Sterol-4-alpha-carboxylate 3-dehydrogenase, decarboxylating GN = Nsdhl	1	Involved in the sequential removal of two C-4 methyl groups in post-squalene cholesterol biosynthesis
O35704	Serine palmitoyltransferase 1 GN = Sptlc1	1	Involved in the pathway sphingolipid metabolism
O55137	Acyl-coenzyme A thioesterase 1 GN = Acot1	1	Catalyze the hydrolysis of acyl-CoAs to the free fatty acid and coenzyme A (CoASH)
Q8BWT1	3-ketoacyl-CoA thiolase, mitochondrial GN = Acaa2	1	Involved in the pathway fatty acid beta oxidation
Q8K0C4	Lanosterol 14-alpha demethylase GN = Cyp51a1	1	Catalyzes C14-demethylation of lanosterol
Q8JZN5	Acyl-CoA dehydrogenase family member 9, mitochondrial GN = Acad9	2	Mitochondrial complex I assembly (By similarity)
Q922J9	Fatty acyl-CoA reductase 1 GN = Far1	1	Catalyzes the reduction of saturated fatty acyl-CoA with chain length C16 or C18 to fatty alcohols
Q99JY0	Trifunctional enzyme subunit beta, mitochondrial GN = Hadhb	1	Involved in the pathway fatty acid beta-oxidation
Q8BLN5	Lanosterol synthase GN = Lss	1	Catalyzes the cyclization of (S)-2,3 oxidosqualene to lanosterol

UniProtID, protein accession number in Swiss-Prot/Treml. ^(a)Protein name given by UniProt database. ^(b)Number of unique peptides matching the protein with a confidence limit of 95%. GN, gene name.

sized in the ER [49]. Recent studies reported that A β PP is also associated with mitochondria in AD affected neurons [50, 51]. This was further supported by the presence of A β PP catabolites (i.e., A β , C99, and AICD) in the mitochondria [9, 52–55]. We report in this study A β PP localization, and processing in MAMs. These results were obtained in two distinct cellular systems (SH-SY5Y and CHO cells) expressing wild type A β PP or harboring two different familial mutations (APP_{swe} and APP_{LDN}) thus demonstrating that the observed presence and processing of WT or mutated A β PP in MAMs is not linked to a specific cell type and not to a specific familial A β PP mutation. This may also reveal that the subcellular localization and proteolytic processing of A β PP in MAMs was not linked to AD familial cases but may also be observed in sporadic cases of AD. These data also suggest that even if trafficking of A β PP may vary according to the mutation, transit through MAMs compartment where A β PP undergo cleavage is a common denominator independent of the nature of the A β PP species.

We also revealed the presence and activity of α -, β -, and γ -secretases in the MAMs. γ -secretase was shown to exert its activity in the plasma membrane but also in other intracellular compartments such as the trans-Golgi network, lysosomes, pure mitochondria fraction, and MAMs [16, 20, 56].

However, it is important to emphasize that even if γ -secretase complex has been found in mitochondria [16], the putative generation of intra-mitochondrial A β and C99 remains puzzling, given the C-terminus-out orientation of A β PP in the outer mitochondrial membrane [9]. A hypothetical model has been proposed stating that A β may be produced at the interface between the ER and mitochondria and then transferred to the mitochondria via the TOM import machinery [9, 44]. Another gap to envision the production of A β in mitochondria was the demonstration of the presence of active β -secretase (BACE-1) in mitochondria or in MAMs. Importantly, we report herein and for the first time the presence of BACE-1 in crude mitochondria and MAMs fractions. Current knowledge on A β PP and BACE-1 trafficking indicates that plasma membrane and endosomes are likely the major sites of BACE-1-mediated cleavage of A β PP [57, 58]. Mature BACE-1 localizes largely within plasma membrane cholesterol-rich lipid rafts [57, 58], and replacing the BACE-1 transmembrane domain with a glycosylphosphatidylinositol anchor extensively targets BACE-1 to lipid rafts and substantially increases A β production [59]. In addition, various types of lipids stimulate BACE-1 activity [60]. Interestingly, a recent study revealed that MAMs behave as a detergent-resistant lipid raft-like domain, consistent with the presence and activity of

β - and γ -secretase in rafts [21, 61–63]. In accordance with the presence of α -CTF fragment in MAMs, we also could measure an *in vitro* α -secretase activity in this compartment in mice brains (Supplementary Figure 5D). Other experiments are still needed to further characterize expression and subcellular distribution and trafficking of α -secretase *in vitro* and *in vivo*. We must take into account that both β - and γ -secretases harbor optimal activities in acidic compartments and that the pH of the ER is almost neutral (pH 7.2), and that of mitochondria matrix is markedly alkaline (pH 8) [64]. Thus, it remains not clear how these enzymes may function in the ER, mitochondria, and MAMs compartments in intact cells. One may postulate that A β PP, or at least its C-terminal fragments (C99/C83), and BACE-1 are transported to the ER-mitochondria contact sites through a mechanism implicating retrograde vesicular trafficking. Indeed, we identified different proteins related to vesicle formation and trafficking in the protein interactome data of MAMs fraction isolated from CHO cells expressing APP_{LDN} (Table 3). Dedicated experiments are necessary to confirm this hypothesis and to further explore the molecular mechanisms underlying BACE-1 localization and *in situ* function in MAMs.

We report an increased ER-mitochondria contact sites in SH-SY5Y cells overexpressing APP_{swe}. Accordingly, an increased ER-mitochondria connectivity was detected in human fibroblasts isolated from individuals with familial AD mutations as well as sporadic AD cases, and in MEFs isolated from mice simple or double knockout for PS1 or PS2 [21]. Similar results were obtained in neuroblastoma cells expressing familial AD-linked PS2 mutations [35]. Interestingly, increased ER-mitochondria connectivity was also reported in a transgenic AD mouse model expressing mutant tau protein [65], and most recently, in cells exposed to apolipoprotein E (ApoE4), a major risk factor for developing sporadic AD [66]. These results and our data may lead to consider MAMs structural alteration as a common denominator underlying the pathogenesis of AD associated to both sporadic and APP/PS-linked familial AD mutations. In agreement with these data, the proteomic approach revealed potential A β PP interactors implicated in the regulation of mitochondrial and ER structure (Dnajc11 and atlastin 3 respectively) (Supplementary Table 1) [67–69]. Accordingly, post-mortem analysis of human AD brain and those of AD mouse models showed altered expression levels of PACS2 and Sigma1R, thus further establishing deregulated MAMs as a hallmark of AD [70]. Moreover,

it is interesting to note that A β PP processing relies on intact ER-mitochondria crosstalk [21]. Thus, cells deficient of MFN2 show diminished MAM function in terms of lipid metabolism and reduced γ -secretase activity by approximately 50% [21]. In turn, exposing hippocampal neurons or neuroblastoma cells to A β enhances ER-mitochondria contacts [70, 71]. These observations are in favor of a vicious cycle where A β PP is processed in MAMs, and its derivative catabolites enhance ER-mitochondria communication and function.

We analyzed a second aspect of MAMs function in AD by combining immunoprecipitation and proteomic mass spectrometric analysis to identify molecules that associate with A β PP and its proteolytic products in MAMs. This approach allows the unbiased and extensive exploration of all possible direct or indirect protein interactors. Nevertheless, we may consider some limitations of the approach used here namely the difficulty to identify interactors of lower expression. This approach failed also to reveal components of γ -secretase and/or β -secretase in MAM proteome analyses. This may be explained by the limitation of immunoprecipitation approach and/or to instable interaction of A β PP and its cleaving enzymes. Although, the strength of our data is supported by the identification of A β PP interactors described in previous studies using distinct experimental approaches (i.e., TIM/TOM complex) [8, 43, 44]. Furthermore, we also revealed new protein partners belonging to cellular pathways known to participate to AD pathogenesis. Interestingly, six out of nine of A β PP and its catabolites interactome related to mitochondrial respiratory chain complexes are components or chaperone of complex I. Previous studies have revealed that A β may cause both a selective defect in complex I activity associated with an increase of intracellular reactive oxygen species [72, 73]. We also unraveled an interaction of A β PP and or its catabolites with the mitochondrial ATP synthase (complex V). Interestingly this enzyme shows reduced activity in AD affected brains [74].

We revealed the interaction of A β PP and its catabolites with several proteins implicated in protein synthesis, maturation, folding, processing, and degradation. Several studies have suggested that altered maturation, processing, and degradation of A β PP holoprotein may drive the availability of A β PP for A β generation [75–78]. It was also suggested that different PPIases regulate A β production, or the toxicity associated with A β [79]. Concerning mitochondria,

we showed that A β PP and its catabolites may directly or indirectly interact with mitochondrial presequence protease (PreP, GN: pitrm), a pitrilysin metalloprotease 1 implicated in the degradation of mitochondrial transit peptides after their cleavage. PreP has been shown to degrade various A β peptides thus clearing mitochondrial (A β) [80, 81]. Accordingly, increased neuronal PreP activity attenuates neuroinflammation and improves mitochondrial and synaptic function in AD mice model [82, 83].

Finally, we delineate in this study the potential interaction of A β PP and its catabolites with different proteins related to fatty-acid metabolism. Accordingly, we show increased neutral lipids accumulation in both neuroblastoma and CHO cells expressing APP_{swe} or APP_{LDN}. Neutral lipids accumulation was reported in different AD study models including PS-mutant MEFs, PS double knockout MEFs, fibroblasts from familial AD patients [21], and more recently fibroblasts exposed to ApoE4 conditioned media [66]. We also revealed that neutral lipids accumulation in our study models is likely linked to A β peptides rather to C99 fragment. A more dedicated study is necessary to investigate the molecular mechanisms underlying lipid droplets accumulation in AD. In particular, lipid droplets are found in close proximity with the ER and mitochondria [37, 38], and the functions of these interactions are still largely unknown. It is, however, noticeable that proteolytic cleavage of A β PP is tightly regulated by several lipids such as cholesterol and sphingolipids. In turn, A β as well as other A β PP processing products play an essential role in regulating lipid homeostasis [84]. Moreover, *in vitro* studies have shown that fatty acids may affect A β secretion [85].

In summary, our data extends the previous studies suggesting MAMs as an important contributor to AD pathogenesis. Maintenance and tight regulation of MAM structure and function could be envisioned as potential therapeutic strategies for AD.

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SUPPLEMENTARY MATERIAL

The supplementary material is available in the electronic version of this article: <http://dx.doi.org/10.3233/JAD-160953>.

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